STIC-Biotech/ChemLib

179011

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F	rom:	
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Steadman, David (AU1652)

Sent:

Wednesday, February 08, 2006 1:18 PM

To:

STIC-Biotech/ChemLib

Subject:

10/601,011 sequence search request

NAME: David Steadman

AU:

1656

Date:

2/8/06

Office:

Remsen 2B05

Mailbox:

Remsen 3C70

Please search the following sequences in commercial databases:

- 1) amino acids 126-388 of SEQ ID NO:1 against amino acid databases.
- 2) amino acids 126-388 of SEQ ID NO:1 against nucleic acid databases.

Please save results to diskette.

Thank you very much.

David J. Steadman, Ph.D. Primary Examiner Art Unit 1656

Protein Crystallography and Recombinant Enzymes

Office: Remsen 2B05 Mailbox: Remsen 3C70 Phone: (571) 272-0942

Type of Search		Vandors and sast w
******		*****
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	. *	

Searcher: _______
Searcher Phone: ______
Date Searcher Picked up: ______
Date completed: ______
Searcher Prep Time: ______
Online Time:

Type of Search

NA#______ AA#:_____

S/L:____ Oligomer:_____

Encode/Transl:_____

Structure #:____ Text:___

Inventor:____ Litigation:

ndors and cost where applicable
STN:
DIALOG:
QUESTEL/ORBIT:
LEXIS/NEXIS:
SEQUENCE SYSTEM:
WWW/Internet:
Other (Specify):

OM protein - protein search, using sw model

February 12, 2006, 18:56:45; Search time 81 Seconds Run on:

(without alignments)

1426.625 Million cell updates/sec

Title: US-10-601-011-1_COPY_126_388

Perfect score: 1383

1 RQWALEDFEIGRPLGKGKFG......SQRPMLREVLEHPWITANSS 263 Sequence:

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2443163 seqs, 439378781 residues

Total number of hits satisfying chosen parameters: 2443163

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : A_Geneseq_21:*

1: geneseqp1980s:*

2: geneseqp1990s:*

3: geneseqp2000s:*

4: geneseqp2001s:*

5: geneseqp2002s:* 6: geneseqp2003as:*

7: geneseqp2003bs:*

8: geneseqp2004s:*

9: geneseqp2005s:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

		8				
Result		Query				
No.	Score	Match	Length	DB	ID	Description
1	1383	100.0	282	9	ADZ47152	Adz47152 Human aur
2	1383	100.0	403	2	AAW18084	Aaw18084 Human Aur
3	1383	100.0	403	2	AAY22476	Aay22476 Human AUR
4	1383	100.0	403	4	AAG67614	Aag67614 Amino aci
5	1383	100.0	403	4	AAG67435	Aag67435 Amino aci
6	1383	100.0	403	6	ABP97367	Abp97367 Human ser
7	1383	100.0	403	6	ABP97366	Abp97366 Human ser
8	1383	100.0	403	6	ABR48160	Abr48160 Human bla
				_		=

-	9	1383	100.0	403	7	ADF61840	•	Adf61840	Human ser	
	10	1383	100.0	403	7	ADN39888			Cancer/an	
	11	1383	100.0	403	8	ADE15337			Truncated	
	12	1383	100.0	403	8	ADH59556			Monoclona	
	13	1383	100.0	403	8	ADK67720			Human mod	
	14	1383	100.0	403	8	ADM72205			Human TAS	
	15	1383	100.0	403	8	ADQ89832			Antagonis	
	16	1383	100.0	403	8	ADR05174		-	Human GTP	
	17	1383	100.0	403	8	ADU06485		Adu06485	Novel bro	
	18	1383	100.0	403	9	ADW15992		Adw15992	Human aur	
	19	1383	100.0	403	9	ADX06369		Adx06369	Cyclin-de	
	20	1383	100.0	403	9	ADY15254		Ady15254	PRO polyp	
	21	1383	100.0	403	9	AEB87732		Aeb87732	Human aur	
	22	1383	100.0	420	5	ABP41958		Abp41958	Human ova	
	23	1383	100.0	461	8	ADX97535		Adx97535	Pancreati	
	24	1377	99.6	300	6	ABP97473		Abp97473	Truncated	
	25	1377	99.6	309	6	ABP97472		Abp97472	Truncated	
	26	1377.	99.6	319	6	ABP97470		Abp97470	Truncated	
	27	1374	99.3	403	7	ADJ31727		Adj31727	Human mit	
	28	1367.5	98.9	402	9	ADZ64605		Adz64605	Kinase do	
	29	1332	96.3	402	7	ABR61579		Abr61579	Human HsA	
	30	1332	96.3	402	7	ADD89969		Add89969	Human can	
	31	1332	96.3	402	8	ADU06504		Adu06504	Novel bro	
	32	1332	96.3	402	9	ADX06367		Adx06367	Cyclin-de	
	33	1332	96.3	402	9	ADY15914		Ady15914	PRO polyp	
	34	1303.5	94.3	320	9	ADY85555		Ady85555	Catalytic	
	35	1284	92.8	403	5	AAO18740		Aao18740	Human NOV	
	36	1174	84.9	279	9	ADZ47153		Adz47153	Clawed fr	
	37	1125	81.3	267	9	ADZ47154		Adz47154	Takifugu	
	38	1047	75.7	280	9	ADZ47158		Adz47158	Clawed fr	
	39	1044	75.5	279	9	ADZ47157			Human aur	
	40	1044	75.5	336	8	ADX97574	,	Adx97574	Pancreati	
	41	1044	75.5	344	2	AAW18083			Human Aur	
	42	1044	75.5	344	2	AAY22475		-	Human AUR	
	43	1044	75.5	344	4	AAG67615		_	Amino aci	
	44	1044	75.5	344	4	AAG67436			Amino aci	
	45	1044	75.5	344	6	ABR48188		Abr48188	Human bla	

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OM protein - protein search, using sw model

Run on: February 12, 2006, 19:03:40; Search time 174 Seconds

(without alignments)

631.547 Million cell updates/sec

Title: US-10-601-011-1_COPY_126_388

Perfect score: 1383

Sequence: 1 RQWALEDFEIGRPLGKGKFG.....SQRPMLREVLEHPWITANSS 263

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 1867569 seqs, 417829326 residues

Total number of hits satisfying chosen parameters: 1867569

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Published Applications AA Main:*

- 1: /cgn2_6/ptodata/1/pubpaa/US07_PUBCOMB.pep:*
- 2: /cgn2_6/ptodata/1/pubpaa/US08_PUBCOMB.pep:*
- 3: /cgn2_6/ptodata/1/pubpaa/US09_PUBCOMB.pep:*
- 4: /cgn2_6/ptodata/1/pubpaa/US10A_PUBCOMB.pep:*
- 5: /cgn2_6/ptodata/1/pubpaa/US10B_PUBCOMB.pep:*
- 6: /cgn2_6/ptodata/1/pubpaa/US11_PUBCOMB.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

	윰				
	Query			•	
Score	Match	Length	DB	ID	Description
1202	100 0	222		119-10-664-421-125	Coguenge 12F Ann
			_		Sequence 125, App
1303	100.0	403	3	US-U9-U12-135A-4	Sequence 4, Appli
1383	100.0	403	3	US-09-784-332-4	Sequence 4, Appli
1383	100.0	403	4	US-10-060-065-12	Sequence 12, Appl
1383	100.0	403	4	US-10-059-585-33	Sequence 33, Appl
1383	100.0	403	4	US-10-209-324-2	Sequence 2, Appli
1383	100.0	403	4	US-10-295-027-1206	Sequence 1206, Ap
1383	100.0	403	4	US-10-087-684-47	Sequence 47, Appl
1383	100.0	403	4	US-10-087-684-48	Sequence 48, Appl
1383	100.0	403	4	US-10-087-684-49	Sequence 49, Appl
1383	100.0	403	4	US-10-218-779-47	Sequence 47, Appl
_	1383 1383 1383 1383 1383 1383 1383 1383	Query Score Match 1383 100.0 1383 100.0 1383 100.0 1383 100.0 1383 100.0 1383 100.0 1383 100.0 1383 100.0 1383 100.0 1383 100.0	Query Score Match Length 1383 100.0 322 1383 100.0 403 1383 100.0 403 1383 100.0 403 1383 100.0 403 1383 100.0 403 1383 100.0 403 1383 100.0 403 1383 100.0 403 1383 100.0 403	Query Score Match Length DB 1383 100.0 322 4 1383 100.0 403 3 1383 100.0 403 4 1383 100.0 403 4 1383 100.0 403 4 1383 100.0 403 4 1383 100.0 403 4 1383 100.0 403 4 1383 100.0 403 4 1383 100.0 403 4 1383 100.0 403 4	Query Score Match Length DB ID 1383 100.0 322 4 US-10-664-421-125 1383 100.0 403 3 US-09-012-135A-4 1383 100.0 403 3 US-09-784-332-4 1383 100.0 403 4 US-10-060-065-12 1383 100.0 403 4 US-10-059-585-33 1383 100.0 403 4 US-10-295-027-1206 1383 100.0 403 4 US-10-295-027-1206 1383 100.0 403 4 US-10-087-684-47 1383 100.0 403 4 US-10-087-684-48 1383 100.0 403 4 US-10-087-684-49

12	1383	100.0	403	4	US-10-218-779-48	Sequence	48, Appl
13	1383	100.0	403	4	US-10-218-779-49	Sequence	49, Appl
14	1383	100.0	403	4	US-10-188-832-35	Sequence	35, Appl
15	1383	100.0	403	5	US-10-751-736-65	Sequence	65, Appl
16	1383	100.0	403	5	US-10-734-126-4	Sequence	4, Appli
17	1383	100.0	403	5	US-10-781-581-187	Sequence	187, App
18	1383	100.0	403	5	US-10-979-375-1	Sequence	1, Appli
19	1383	100.0	403	5	US-10-745-237-262	Sequence	262, App
20	1383	100.0	403	5	US-10-965-313-1	Sequence	1, Appli
21	1383	100.0	403	6	US-11-037-713-2	Sequence	2, Appli
22	1383	100.0	403	6	US-11-092-809-1	Sequence	1, Appli
23	1383	100.0	403	6	US-11-092-863-1	Sequence	1, Appli
24	1383	100.0	420	4	US-10-264-049-3090	Sequence	3090, Ap
25	1374	99.3	403	4	US-10-026-021-7	Sequence	7, Appli
26	1374	99.3	403	4	US-10-087-684-50	Sequence	50, Appl
27	1374	99.3	403	4	US-10-218-779-50		50, Appl
28	1332	96.3	321	5	US-10-941-635-127	Sequence	127, App
29	1332	96.3	402	4	US-10-087-684-51	Sequence	51, Appl
30	1332	96.3	402	4	US-10-218-779-51	Sequence	51, Appl
31	1332	96.3	402	5	US-10-509-599-14	Sequence	14, Appl
32	1284	92.8	403	4	US-10-087-684-14	Sequence	14, Appl
33	1284	92.8	403	4	US-10-218-779-14	Sequence	14, Appl
34	1044	75.5	344	3	US-09-012-135A-3	Sequence	3, Appli
35	1044	75.5	344	3	US-09-784-332-3	Sequence	3, Appli
36	1044	75.5	344	4	US-10-060-065-13	Sequence	13, Appl
37	1044	75.5	344	4	US-10-059-585-34	Sequence	34, Appl
38	1044	75.5	344	4	US-10-171-311-214	Sequence	214, App
39	1044	75.5	344	4	US-10-295-027-203		203, App
40	1044	75.5	344	4	US-10-173-999-101	Sequence	101, App
41	1044	75.5	344	4	US-10-188-832-93	Sequence	93, Appl
42	1044	75.5	344	5	US-10-734-126-3	Sequence	3, Appli
43	1044	75.5	344	5	US-10-745-237-264	Sequence	264, App
44	1044	75.5	344	5	US-10-965-313-2	Sequence	2, Appli
45	1044	75.5	344	6	US-11-037-713-30	Sequence	30, Appl

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OM protein - nucleic search, using frame plus p2n model

Run on: February 14, 2006, 20:29:25; Search time 3993 Seconds

(without alignments)

3744.009 Million cell updates/sec

Title: US-10-601-011-1 COPY 126 388

Perfect score: 1383

Sequence: 1 RQWALEDFEIGRPLGKGKFG.....SQRPMLREVLEHPWITANSS 263

Scoring table: BLOSUM62

Xgapop 10.0 , Xgapext Ygapop 10.0 , Ygapext 0.5 Fgapop 6.0 , Fgapext 7.0 6.0 , Delext Delop 7.0

Searched: 5883141 seqs, 28421725653 residues

Total number of hits satisfying chosen parameters: 11766282

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Command line parameters:

- -MODEL=frame+_p2n.model -DEV=xlp
- -Q=/abss/ABSSWEB_spool/US10601011/runat_12022006_150911_13333/app_query.fasta_1
- -DB=GenEmbl -QFMT=fastap -SUFFIX=rge -MINMATCH=0.1 -LOOPCL=0 -LOOPEXT=0
- -UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi -LIST=45
- -DOCALIGN=200 -THR_SCORE=pct -THR_MAX=100 -THR_MIN=0 -ALIGN=15 -MODE=LOCAL
- -OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=200000000 -HOST=abss05p
- -USER=US10601011_@CGN_1_1_7415_@runat_12022006_150911_13333 -NCPU=6 -ICPU=3 -NO_MMAP -NEG_SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG -DEV_TIMEOUT=120
- -WARN TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6 -FGAPEXT=7
- -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7

Database : GenEmbl: *

- 1: gb ba:*
- 2: gb_in:*
- 3: gb env:*
- 4: gb om:*
- 5: gb ov:*
- 6: gb_pat:*
- 7: gb_ph:*
- 8: gb_pr:*
- 9: gb_ro:* 10: gb_sts:*
- 11: gb_sy:*
- 12: gb_un:*

13: gb_vi:* 14: gb_htg:* 15: gb_pl:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Res	ult		Query				
	No.	Score	Match	Length	DB	ID	Description
	1	1383	100.0	1212	8	AF008551	AF008551 Homo sapi
	2	1383	100.0	1212	11	AY335576	AY335576 Synthetic
	3	1383	100.0	1212	11	AY892410	AY892410 Synthetic
	4	1383	100.0	1212	11	BT008056	BT008056 Synthetic
	5	1383	100.0	2024	8	BC006423	BC006423 Homo sapi
С	6	1383	100.0	2064	10	BV177260	BV177260 sqnm93669
•	7	1383	100.0	2120	8	BC002499	BC002499 Homo sapi
	8	1383	100.0	2121	6	CS119570	CS119570 Sequence
	9	1383	100.0	2128	8	BC001280	BC001280 Homo sapi
	10	1383	100.0	2135	6	CS119568	CS119568 Sequence
	11	1383	100.0	2169	8	BC027464	BC027464 Homo sapi
	12	1383	100.0	2198	6	AR078203	AR078203 Sequence
	13	1383	100.0	2198	6	AR081975	AR081975 Sequence
	14	1383	100.0	2198	6	AR139681	AR139681 Sequence
	15	1383	100.0	2198	6	BD140128	BD140128 Diagnosis
	16	1383	100.0	2198	6	AR491644	AR491644 Sequence
	17	1383	100.0	2198	6	AR630705	AR630705 Sequence
	18	1383	100.0	2231	6	CS119569	CS119569 Sequence
	19	1383	100.0	2237	6	CQ728025	CQ728025 Sequence
	20	1383	100.0	2237	6	AX281843	AX281843 Sequence
	21	1383	100.0	2245	6	CS119567	CS119567 Sequence
	22	1383	100.0	2253	6	CQ759659	CQ759659 Sequence
	23	1383	100.0	2253	6	CQ894702	CQ894702 Sequence
	24	1383	100.0	2253	6	CQ981365	CQ981365 Sequence
	25	1383	100.0	2253	6	CS031553	CS031553 Sequence
	26	1383	100.0	2253	6	CS040505	CS040505 Sequence
	27	1383	100.0	2253	6	CS118077	CS118077 Sequence
	28	1383	100.0	2253	6	AR198495	AR198495 Sequence
	29	1383	100.0	2253	8	AF011468	AF011468 Homo sapi
С	30	1383	100.0	2284	6	CQ414603	CQ414603 Sequence
	31	1383	100.0	2346	6	CS119565	CS119565 Sequence
	32	1383	100.0	2347	6	AR648112	AR648112 Sequence
	33	1383	100.0	2554	6	CS119566	CS119566 Sequence
	34	1363	98.6	2148	6	AX281842	AX281842 Sequence
	35	1349	97.5	1934	8	AB168134	AB168134 Macaca fa
	36	1334.5	96.5	1218	4	AB196773	AB196773 Sus scrof
	37	1332	96.3	1209	6	CQ847956	CQ847956 Sequence
	38	1332	96.3	1209	6	AX934319	AX934319 Sequence
	39	1332	96.3	2033	6	CQ981384	CQ981384 Sequence
	40	1332	96.3	2033	6	CS032213	CS032213 Sequence
	41	1332	96.3	2033	6	CS041165	CS041165 Sequence
	42	1332	96.3	2033	6	AR198502	AR198502 Sequence
	43	1332	96.3	2033	6	AX925606	AX925606 Sequence
	44	1332	96.3	2033	8	D84212	D84212 Homo sapien

OM protein - nucleic search, using frame plus p2n model

Run on: February 14, 2006, 21:16:44; Search time 187 Seconds

(without alignments)

2499.993 Million cell updates/sec

Title: US-10-601-011-1 COPY 126 388

Perfect score: 1383

Sequence: 1 RQWALEDFEIGRPLGKGKFG.....SQRPMLREVLEHPWITANSS 263

Scoring table: BLOSUM62

Xgapop 10.0 , Xgapext 0.5 Ygapop 10.0 , Ygapext 0.5 Fgapop 6.0 , Fgapext 7.0 Delop 6.0 , Delext 7.0

Searched: 1303057 seqs, 888780828 residues

Total number of hits satisfying chosen parameters: 2606114

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Command line parameters:

- -MODEL=frame+_p2n.model -DEV=xlp
- -Q=/abss/ABSSWEB_spool/US10601011/runat_12022006_150916_13432/app_query.fasta_1
- -DB=Issued_Patents_NA -QFMT=fastap -SUFFIX=rni -MINMATCH=0.1 -LOOPCL=0
- -LOOPEXT=0 -UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi
- -LIST=45 -DOCALIGN=200 -THR_SCORE=pct -THR_MAX=100 -THR MIN=0 -ALIGN=15
- -MODE=LOCAL -OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=2000000000
- -HOST=abss06p -USER=US10601011_@CGN_1_1_290_@runat_12022006_150916_13432
- -NCPU=6 -ICPU=3 -NO_MMAP -NEG_SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG
- -DEV_TIMEOUT=120 -WARN TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6
- -FGAPEXT=7 -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7

Database : Issued_Patents_NA:*

- 1: /cgn2_6/ptodata/1/ina/1_COMB.seq:*
- 2: /cgn2_6/ptodata/1/ina/5 COMB.seq:*
- 3: /cgn2_6/ptodata/1/ina/6A_COMB.seq:*
- 4: /cgn2_6/ptodata/1/ina/6B_COMB.seq:*
- 5: /cgn2_6/ptodata/1/ina/H COMB.seq:*
- 6: /cgn2_6/ptodata/1/ina/PCTUS_COMB.seq:*
- 7: /cgn2_6/ptodata/1/ina/PP COMB.seq:*
- 8: /cgn2_6/ptodata/1/ina/RE_COMB.seq:*
- 9: /cgn2_6/ptodata/1/ina/backfiles1.seq:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed,

and is derived by analysis of the total score distribution.

		&			SUMMARIES	
Pocult						
Result No.	Score	Query	Length	מת	ID	Doggrintion
NO.	Score	Macch	nengun	םע	10	Description
1	1383	100.0	2198	2	US-08-755-728-2	Sequence 2, Appli
2	1383	100.0	2198	2	US-08-974-655-2	Sequence 2, Appli
3	1383	100.0	2198	3	US-09-283-011-2	Sequence 2, Appli
4	1383	100.0	2198	3	US-09-012-135A-2	Sequence 2, Appli
5	1383	100.0	2198	3	US-09-784-332-2	Sequence 2, Appli
6	1383	100.0	2237	3	US-09-949-016-4590	Sequence 4590, Ap
7	1383	100.0	2237	3	US-09-949-016-4591	Sequence 4591, Ap
8	1383	100.0	2253	3	US-09-660-925B-3	Sequence 3, Appli
9	1383	100.0	2347	3	US-10-209-324-1	Sequence 1, Appli
10	1332	96.3	2033	3	US-09-660-925B-10	Sequence 10, Appl
11	1044	75.5	1224	3	US-09-949-016-3955	Sequence 3955, Ap
12	1044	75.5	1244	2	US-08-755-728-1	Sequence 1, Appli
13	1044	75.5	1244	2	US-08-974-655-1	Sequence 1, Appli
14	1044	75.5	1244	3	US-09-283-011-1	Sequence 1, Appli
15	1044	75.5	1244	3	US-09-012-135A-1	Sequence 1, Appli
16	1044	75.5	1244	3	US-09-784-332-1	Sequence 1, Appli
17	1032.5	74.7	1281	2	US-09-016-000-7	Sequence 7, Appli
18	1029	74.4	1815	3	US-09-485-534-3	Sequence 3, Appli
19	685	49.5	9822	3	US-09-949-016-15697	Sequence 15697, A
20	666.5	48.2	1533	3	US-09-248-796A-4410	Sequence 4410, Ap
21	528	38.2	747	3	US-09-270-767-15286	Sequence 15286, A
22	527	38.1	26867	3	US-09-949-016-16332	Sequence 16332, A
23	527	38.1	26867	3	US-09-949-016-16333	Sequence 16333, A
24	505.5	36.6	3937	3	US-09-620-312D-280	Sequence 280, App
25	501.5	36.3	1453	2	US-08-252-995D-1	Sequence 1, Appli
26	501.5	36.3	1453	2	US-08-834-108-1	Sequence 1, Appli
27	501.5	36.3	1600	2	US-08-252-995D-5	Sequence 5, Appli
28	501.5	36.3	1600	2	US-08-834-108-5	Sequence 5, Appli
29	501.5	36.3	3447	2	US-08-252-995D-3	Sequence 3, Appli
30	501.5	36.3	3447	2	US-08-834-108-3	Sequence 3, Appli
31	480.5	34.7	781	3	US-09-533-559-7482	Sequence 7482, Ap
32	468	33.8	1194	3	US-09-487-558B-399	Sequence 399, App
33	466	33.7	1197	3	US-09-487-558B-403	Sequence 403, App
34	452	32.7	1929	3	US-09-359-161-4	Sequence 4, Appli
35	444	32.1	3857	3	US-10-464-939-1	Sequence 1, Appli
36	444	32.1	4790	3	US-10-464-939-3	Sequence 3, Appli
37	443	32.0	1143	3	US-09-487-558B-401	Sequence 401, App
38	437.5	31.6	993	3	US-09-248-796A-4394	Sequence 4394, Ap
39	435.5	31.5	1732	3	US-09-430-564-1	Sequence 1, Appli
40	435.5	31.5	1816	3	US-09-762-258-1	Sequence 1, Appli
41	433.5	31.3	2698	2	US-08-677-298-1	Sequence 1, Appli
42	433.5	31.3	2912	3	US-09-949-016-2363	Sequence 2363, Ap
43	433.5	31.3	2914	3	US-09-949-016-343	Sequence 343, App
44	432	31.2	1212	3	US-09-248-796A-6416	Sequence 6416, Ap
45	430	31.1	1539	3	US-09-633-328B-1	Sequence 1, Appli

OM protein - nucleic search, using frame plus p2n model

Run on: February 14, 2006, 21:21:44; Search time 1404 Seconds

(without alignments)

168.433 Million cell updates/sec

Title: US-10-601-011-1 COPY 126 388

Perfect score: 1383

Sequence: 1 RQWALEDFEIGRPLGKGKFG......SQRPMLREVLEHPWITANSS 263

Scoring table: BLOSUM62

Xgapop 10.0 , Xgapext 0.5 Ygapop 10.0 , Ygapext 0.5 Fgapop 6.0 , Fgapext 7.0 Delop 6.0 , Delext 7.0

Searched: 6240305 seqs, 449581930 residues

Total number of hits satisfying chosen parameters: 12480610

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Command line parameters:

- -MODEL=frame+_p2n.model -DEV=xlp
- -Q=/abss/ABSSWEB spool/US10601011/runat 12022006 150920 13536/app query.fasta 1
- -DB=Published Applications NA_New -QFMT=fastap -SUFFIX=rnpbn -MINMATCH=0.1
- -LOOPCL=0 -LOOPEXT=0 -UNITS=bits -START=1 -END=-1 -MATRIX=blosum62
- -TRANS=human40.cdi -LIST=45 -DOCALIGN=200 -THR SCORE=pct -THR MAX=100
- -THR_MIN=0 -ALIGN=15 -MODE=LOCAL -OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0
- -MAXLEN=2000000000 -HOST=abss06p
- -USER=US10601011_@CGN_1_1_335_@runat_12022006_150920_13536 -NCPU=6 -ICPU=3
- -NO MMAP -NEG SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG -DEV TIMEOUT=120
- -WARN TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6 -FGAPEXT=7
- -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7

Published Applications NA New:* Database :

- 1: /cgn2 6/ptodata/1/pubpna/US08 NEW PUB.seq:*
- 2: /cgn2 6/ptodata/1/pubpna/US06 NEW PUB.seq:*
- /cgn2 6/ptodata/1/pubpna/US07 NEW PUB.seq:*
- 4: /cgn2_6/ptodata/1/pubpna/PCT_NEW_PUB.seq:*
- 5: /cgn2_6/ptodata/1/pubpna/US09_NEW_PUB.seq:* 6:
- /cgn2_6/ptodata/1/pubpna/US10_NEW_PUB.seq:* 7:
- /cgn2_6/ptodata/1/pubpna/US10_NEW_PUB.seq1:*
- 8: /cgn2_6/ptodata/1/pubpna/US11 NEW PUB.seq:*
- 9: /cgn2_6/ptodata/1/pubpna/US11_NEW_PUB.seq2:*
- 10: /cgn2 6/ptodata/1/pubpna/US11 NEW PUB.seg3:*
- 11: /cgn2 6/ptodata/1/pubpna/US11 NEW PUB.seq4:*

12: /cgn2_6/ptodata/1/pubpna/US60_NEW_PUB.seq:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

			8				
Res	ult		Query				
	No.	Score	Match	Length	DB	ID	Description
	1	1383	100.0	2253	7	US-10-770-726-39	Sequence 39, Appl
	2	1383	100.0	2253	11	US-11-000-688-124	Sequence 124, App
	3	1383	100.0	2347	7	US-10-770-726-41	Sequence 41, Appl
	4	1332	96.3	2033	7	US-10-523-477-4	Sequence 4, Appli
	5	1044	75.5	1224	7	US-10-523-477-2	Sequence 2, Appli
	6	1044	75.5	1224	7	US-10-770-726-40	Sequence 40, Appl
	7	1044	75.5	1224	11	US-11-136-815A-4	Sequence 4, Appli
	8	1004	72.6	1327	7	US-10-523-477-3	Sequence 3, Appli
	9	508.5	36.8	3331	7	US-10-770-726-42	Sequence 42, Appl
	10	435.5	31.5	1680	7	US-10-770-726-37	Sequence 37, Appl
	11	433.5	31.3	3394	7	US-10-995-561-364	Sequence 364, App
	12	433.5	31.3	3442	7	US-10-995-561-361	Sequence 361, App
	13	433.5	31.3	3466	7	US-10-995-561-363	Sequence 363, App
	14	433.5	31.3	3487	7	US-10-995-561-359	Sequence 359, App
	15	433.5	31.3	3514	7	US-10-995-561-360	Sequence 360, App
	16	428	30.9	1891	11	US-11-000-688-55	Sequence 55, Appl
	17	421.5	30.5	1800	11	US-11-136-527-3715	Sequence 3715, Ap
	18	421.5	30.5	2301	11	US-11-136-527-3070	Sequence 3070, Ap
	19	421.5	30.5	2346	11	US-11-117-013-3	Sequence 3, Appli
	20	421.5	30.5	2346	11	US-11-000-688-1538	Sequence 1538, Ap
	21	421	30.4	2972	11	US-11-169-041-72	Sequence 72, Appl
	22	419.5	30.3	2572	11	US-11-136-527-2589	Sequence 2589, Ap
	23	419	30.3	3511	7	US-10-995-561-362	Sequence 362, App
	24	415.5	30.0	2301	11	US-11-151-601-9	Sequence 9, Appli
	25	415.5	30.0	3552	11	US-11-151-601-7	Sequence 7, Appli
	26	414	29.9	2213	7	US-10-770-726-31	Sequence 31, Appl
	27	409	29.6	2190	7	US-10-947-249-104	Sequence 104, App
	28	405.5	29.3	2615	11	US-11-136-527-2878	Sequence 2878, Ap
	29	404	29.2	1915	11	US-11-136-527-2876	Sequence 2876, Ap
	30	403	29.1	1400	11	US-11-136-527-7767	Sequence 7767, Ap
	31	403	29.1	1402	11	US-11-136-527-3671	Sequence 3671, Ap
	32	403	29.1	1442	11	US-11-136-527-1916	Sequence 1916, Ap
	33	402	29.1	3144	11	US-11-136-527-2887	Sequence 2887, Ap
	34	401.5	29.0	1715	7	US-10-947-249-56	Sequence 56, Appl
С	35	401.5	29.0	3206	7	US-10-821-234-306	Sequence 306, App
	36	401	29.0	2226	11	US-11-136-527-3162	Sequence 3162, Ap
	37	401	29.0	4278	11	US-11-128-061-532	Sequence 532, App
	38	401	29.0	4278	11	US-11-128-049-532	Sequence 532, App
	39	400	28.9	2902	11	US-11-241-056-4	Sequence 4, Appli
C	40	400	28.9	35893	7	US-10-860-436-2	Sequence 2, Appli
	41	399	28.9	1122	11	US-11-136-527-1805	Sequence 1805, Ap
	42	399	28.9	2685	11	US-11-000-688-270	Sequence 270, App
	43	398.5	28.8	1618	11	US-11-136-527-2397	Sequence 2397, Ap
	44	397	28.7	2312	11	US-11-136-527-3389	Sequence 3389, Ap
	45	396.5	28.7	2349	11	US-11-136-527-2338	Sequence 2338, Ap
						== ===	1 up

OM protein - nucleic search, using frame plus p2n model Run on: February 14, 2006, 20:38:26; Search time 3735 Seconds (without alignments) 3294.512 Million cell updates/sec US-10-601-011-1_COPY_126_388 Perfect score: 1383 1 RQWALEDFEIGRPLGKGKFG......SQRPMLREVLEHPWITANSS 263 Sequence: Scoring table: BLOSUM62 Xgapop 10.0 , Xgapext 0.5 Ygapop 10.0 , Ygapext 0.5 Fgapop 6.0 , Fgapext 7.0 6.0 , Delext Delop 7.0 Searched: 41078325 seqs, 23393541228 residues Total number of hits satisfying chosen parameters: 82156650 Minimum DB seq length: 0 Maximum DB seq length: 2000000000 Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries Command line parameters: -MODEL=frame+ p2n.model -DEV=xlp -Q=/abss/ABSSWEB_spool/US10601011/runat_12022006_150913_13375/app_query.fasta_1 -DB=EST -QFMT=fastap -SUFFIX=rst -MINMATCH=0.1 -LOOPCL=0 -LOOPEXT=0 -UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi -LIST=45 -DOCALIGN=200 -THR_SCORE=pct -THR_MAX=100 -THR_MIN=0 -ALIGN=15 -MODE=LOCAL -OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=2000000000 -HOST=abss05p -USER=US10601011_@CGN_1_1_8010_@runat_12022006_150913_13375 -NCPU=6 -ICPU=3 -NO MMAP -NEG SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG -DEV_TIMEOUT=120 -WARN_TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6 -FGAPEXT=7 -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7 Database : EST:* 1: gb_est1:* 2: gb_est2:* 3: gb_est3:* 4: gb htc:* 5: gb est4:*

Pred. No. is the number of results predicted by chance to have a

6: gb_est5:*
7: gb_est6:*
8: gb_est7:*
9: gb_gss1:*
10: gb_gss2:*
11: gb_gss3:*

score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

			•			SUMMARIES	
_			*				
Res		_	Query				
	No.	Score	Match	Length	DB	ID	Description
		1202	100 0				20050151 77
	1	1383	100.0	1212		DQ050151	DQ050151 Homo sapi
	2	1383	100.0	1841	4	CR616247	CR616247 full-leng
	3	1383	100.0	1992	4	CR621289	CR621289 full-leng
	4	1383	100.0	2025	4	CR611278	CR611278 full-leng
	5	1365	98.7	793	8	CX782148	CX782148 HESC3_12_
	6	1322	95.6	811	1	AU124034	AU124034 AU124034
С	7	1320.5	95.5	930	5	BX372079	BX372079 BX372079
	8	1316	95.2	1626	4	BC086984	BC086984 Rattus no
	9	1309	94.6	961	5	BQ685975	BQ685975 AGENCOURT
	10	1305	94.4	1843	4	AK085861	AK085861 Mus muscu
	11	1294	93.6	1852	4	AK077513	AK077513 Mus muscu
	12	1274	92.1	855	6	CA488875	CA488875 AGENCOURT
	13	1264	91.4	1923	4	BC050630	BC050630 Homo sapi
	14	1208	87.3	720	7	CN273685	CN273685 170006008
	15	1177	85.1	854	8	CX747442	CX747442 JGI_ANHP4
	16	1172	84.7	834	8	CX873538	CX873538 HESC4_82_
	17	1166	84.3	696	7	CN791002	CN791002 4125663 B
	18	1158	83.7	883	6	CA971494	CA971494 AGENCOURT
	19	1131.5	81.8	1203	5	BU510028	BU510028 AGENCOURT
	20	1118.5	80.9	850	7	CK023559	CK023559 AGENCOURT
	21	1118	80.8	936	5	BU915671	BU915671 AGENCOURT
	22	1117	80.8	838	8	CX219837	CX219837 MNS35076
	23	1100.5	79.6	1055	2	BG027828	BG027828 602294784
	24	1100	79.5	736	5	BU210010	BU210010 602906028
	25	1090	78.8	1010	2	BG027837	BG027837 602294795
	26	1087	78.6	635	2	BE299015	BE299015 601119568
	27	1087	78.6	730	1	AW957623	AW957623 EST369693
	28	1045	75.6	911	5	BU111086	BU111086 603127273
	29	1043	75.4	978	4	CR591471	CR591471 full-leng
	30	1043	75.4	1162	4	CR604968	CR604968 full-leng
	31	1043	75.4	1199	4	CR600836	CR600836 full-leng
	32	1043	75.4	1199	4	CR603941	CR603941 full-leng
	33	1043	75.4	1227	4	CR605805	CR605805 full-leng
	34	1042.5	75.4	859	2	BG290257	BG290257 602384959
	35	1036	74.9	731	7	CR370688	CR370688 CR370688
	36	1036	74.9	747	5	BX304728	BX304728 BX304728
	37	1031	74.5	1923	4	AK075951	AK075951 Mus muscu
	38	1020	73.8	607	2	BG077290	BG077290 H3013F02-
	39	1017	73.5	826	10	AY421575	AY421575 Homo sapi
	40	1016	73.5	810	8	DN510979	DN510979 HL02021B1
	41	1015.5	73.4	1065	5	BX425271	BX425271 BX425271
	42	1013.5	73.3	890	5	BU105061	BU105061 603007034
	43	1011	73.1	890	8	DN110878	DN110878 1109950 M
	44	1007	72.8	674	5	BX846885	BX846885 BX846885
	45	1004	72.6	784	8	CX232498	CX232498 MBM12224
				_			

OM protein - protein search, using sw model

Run on: February 12, 2006, 19:02:55; Search time 47 Seconds

(without alignments)

462.632 Million cell updates/sec

Title: US-10-601-011-1 COPY 126 388

Perfect score: 1383

Sequence: 1 RQWALEDFEIGRPLGKGKFG.....SQRPMLREVLEHPWITANSS 263

Scoring table: BLOSUM62

Gapop 10.0, Gapext 0.5

Searched: 572060 seqs, 82675679 residues

Total number of hits satisfying chosen parameters: 572060

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Issued Patents AA:*

1: /cgn2_6/ptodata/1/iaa/5_COMB.pep:*

2: /cgn2_6/ptodata/1/iaa/6 COMB.pep:*

3: /cgn2_6/ptodata/1/iaa/H_COMB.pep:*

4: /cgn2_6/ptodata/1/iaa/PCTUS_COMB.pep:*

5: /cgn2_6/ptodata/1/iaa/RE_COMB.pep:*

6: /cgn2_6/ptodata/1/iaa/backfiles1.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

		€				
Result		Query				
No.	Score	Match	Length	DB	ID	Description
1	1383	100.0	403	1	US-08-755-728-4	Sequence 4, Appli
2	1383	100.0	403	1	US-08-974-655-4	Sequence 4, Appli
3	1383	100.0	403	2.	US-09-283-011-4	Sequence 4, Appli
4	1383	100.0	403	2	US-09-012-135A-4	Sequence 4, Appli
5	1383	100.0	403	2	US-09-784-332-4	Sequence 4, Appli
6	1383	100.0	403	2	US-10-209-324-2	Sequence 2, Appli
7	1383	100.0	488	2	US-09-949-016-10461	Sequence 10461, A
8	1383	100.0	488	2	US-09-949-016-10462	Sequence 10462, A
9	1044	75.5	344	1	US-08-755-728-3	Sequence 3, Appli
10	1044	75.5	344	1	US-08-974-655-3	Sequence 3, Appli
11	1044	75.5	344	2	US-09-283-011-3	Sequence 3, Appli

12	1044	75.5	344	2	US-09-012-135A-3	Sequence 3, Appli
13	1044	75.5	344	2	US-09-784-332-3	Sequence 3, Appli
14	1044	75.5	363	2	US-09-949-016-9826	Sequence 9826, Ap
15	1032.5	74.7	347	1	US-09-016-000-1	Sequence 1, Appli
16	1029	74.4	343	2	US-09-485-534-4	Sequence 4, Appli
17	891	64.4	436	2	US-09-485-534-18	Sequence 18, Appl
18	701.5	50.7	412	2	US-09-485-534-19	Sequence 19, Appl
19	666.5	48.2	510	2	US-09-248-796A-18513	Sequence 18513, A
20	501.5	36.3	273	1	US-08-252-995D-10	Sequence 10, Appl
21	501.5	36.3	273	1	US-08-834-108-10	Sequence 10, Appl
22	501.5	36.3	416	1	US-08-252-995D-2	Sequence 2, Appli
23	501.5	36.3	416	1	US-08-834-108-2	Sequence 2, Appli
24	501.5	36.3	464	1	US-08-252-995D-6	Sequence 6, Appli
25	501.5	36.3	464	1	US-08-834-108-6	Sequence 6, Appli
26	501.5	36.3	925	1	US-08-252-995D-4	. Sequence 4, Appli
27	501.5	36.3	925	1	US-08-834-108-4	Sequence 4, Appli
28	497.5	36.0	256	2	US-09-964-956-72	Sequence 72, Appl
29	481.5	34.8	256	2	US-09-964-956-74	Sequence 74, Appl
30	474	34.3	260	1	US-07-857-224B-6	Sequence 6, Appli
31	468	33.8	397	2	US-09-487-558B-400	Sequence 400, App
32	466	33.7	398	2	US-09-538-092-501	Sequence 501, App
33	466	33.7	398	2	US-09-487-558B-404	Sequence 404, App
34	457	33.0	227	2	US-09-270-767-46850	Sequence 46850, A
35	453.5	32.8	446	2	US-09-824-735-2	Sequence 2, Appli
36	453	32.8	260	1	US-07-857-224B-8	Sequence 8, Appli
37	447	32.3	258	1	US-07-857-224B-5	Sequence 5, Appli
38	445	32.2	511	2	US-09-633-328B-4	Sequence 4, Appli
39	444	32.1	832	2	US-10-464-939-2	Sequence 2, Appli
40	444	32.1	1058	2	US-10-464-939-4	Sequence 4, Appli
41	443	32.0	380	2	US-09-487-558B-402	Sequence 402, App
42	437.5	31.6	330	2	US-09 - 248-796A-18497	Sequence 18497, A
43	437	31.6	260	1	US-07-857-224B-7	Sequence 7, Appli
44	435.5	31.5	268	2	US-09-430-564-12	Sequence 12, Appl
45	435.5	31.5	482	2	US-09-430-564-2	Sequence 2, Appli

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OM protein - protein search, using sw model

Run on: February 12, 2006, 19:04:35; Search time 16 Seconds

(without alignments)

215.702 Million cell updates/sec

Title: US-10-601-011-1 COPY 126 388

Perfect score: 1383

Sequence: 1 RQWALEDFEIGRPLGKGKFG.....SQRPMLREVLEHPWITANSS 263

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 97014 seqs, 13122538 residues

Total number of hits satisfying chosen parameters: 97014

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Published Applications AA New:*

1: /cgn2 6/ptodata/1/pubpaa/US08 NEW PUB.pep:*

2: /cgn2_6/ptodata/1/pubpaa/US06_NEW_PUB.pep:*

3: /cgn2_6/ptodata/1/pubpaa/US07 NEW PUB.pep:*

4: /cgn2_6/ptodata/1/pubpaa/PCT NEW PUB.pep:*

5: /cgn2_6/ptodata/1/pubpaa/US09 NEW PUB.pep:*

6: /cgn2_6/ptodata/1/pubpaa/US10_NEW_PUB.pep:*

7: /cgn2_6/ptodata/1/pubpaa/US11_NEW_PUB.pep:*

8: /cgn2_6/ptodata/1/pubpaa/US60 NEW PUB.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

		ક				
Result		Query				
No.	Score	Match	Length	DB	ID	Description
1	1383	100.0	403	 6	US-10-523-477-11	Sequence 11, Appl
2	1383	100.0	403	6	US-10-770-726-83	Sequence 83, Appl
3	1383	100.0	403	6	US-10-770-726-85	Sequence 85, Appl
4	1383	100.0	403	7	US-11-109-156-12	Sequence 12, Appl
5	1383	100.0	403	7	US-11-092-168-1	Sequence 1, Appli
6	1044	75.5	344	6	US-10-523-477-9	Sequence 9, Appli
7	1044	75.5	344	6	US-10-770-726-84	Sequence 84, Appl
8	1044	75.5	344	7	US-11-109-156-13	Sequence 13, Appl
9	1044	75.5	344	7	US-11-092-168-2	Sequence 2, Appli

10	1004	72.6	275	6	US-10-523-477-10	Sequence 10, Appl
11	508.5	36.8	970	6	US-10-770-726-86	Sequence 86, Appl
12	497.5	36.0	256	6	US-10-877-346-72	Sequence 72, Appl
13	497.5	36.0	256	7	US-11-113-424-183	Sequence 183, App
14	481.5	34.8	256	6	US-10-877-346-74	Sequence 74, Appl
15	466	33.7	398	7	US-11-132-142-7	Sequence 7, Appli
16	445	32.2	381	7	US-11-132-142-9	Sequence 9, Appli
17	435.5	31.5	480	7	US-11-132-142-6	Sequence 6, Appli
18	435.5	31.5	495	6	US-10-770-726-81	Sequence 81, Appl
19	434	31.4	515	7	US-11-132-142-8	Sequence 8, Appli
20	433.5	31.3	713	6	US-10-995-561-881	Sequence 881, App
21	433.5	31.3	729	6	US-10-995-561-878	Sequence 878, App
22	433.5	31.3	737	6	US-10-995-561-880	Sequence 880, App
23	433.5	31.3	744	6	US-10-995-561-876	Sequence 876, App
24	433.5	31.3	753	6	US-10-995-561-877	Sequence 877, App
25	423	30.6	353	7	US-11-132-142-11	Sequence 11, Appl
26	421	30.4	685	7	US-11-169-041-194	Sequence 194, App
27	419	30.3	752	6	US-10-995-561-879	Sequence 879, App
28	416	30.1	462	7	US-11-132-142-10	Sequence 10, Appl
29	415.5	30.0	766	7	US-11-151-601-8	Sequence 8, Appli
30	414	29.9	603	6	US-10-770-726-75	Sequence 75, Appl
31	412	29.8	277	7	US-11-151-601-4	Sequence 4, Appli
32	412	29.8	351	7	US-11-132-142-13	Sequence 13, Appl
33	411	29.7	548	7	US-11-132-142-5	Sequence 5, Appli
34	410.5	29.7	278	6	US-10-055-877-149	Sequence 149, App
35	410.5	29.7	278	7	US-11-103-065-4	Sequence 4, Appli
36	410.5	29.7	278	7	US-11-151-601-10	Sequence 10, Appl
37	409	29.6	740	6	US-10-878-556A-129	Sequence 129, App
38	401.5	29.0	543	6	US-10-821-234-1158	Sequence 1158, Ap
39	401	29.0	343	7	US-11-092-168-3	Sequence 3, Appli
40	400	28.9	631	7	US-11-241-056-11	Sequence 11, Appl
41	399	28.9	332	6	US-10-497-767-3	Sequence 3, Appli
42	399	28.9	351	7	US-11-132-142-14	Sequence 14, Appl
43	396	28.6	395	7	US-11-132-142-12	Sequence 12, Appl
44	393	28.4	350	6	US-10-497-767-4	Sequence 4, Appli
45	390	28.2	514	7	US-11-125-295-2	Sequence 2, Appli

OM protein - nucleic search, using frame plus p2n model

Run on: February 14, 2006, 20:22:30; Search time 517 Seconds

(without alignments)

3390.356 Million cell updates/sec

Title: US-10-601-011-1 COPY 126 388

Perfect score: 1383

Sequence: 1 RQWALEDFEIGRPLGKGKFG.....SQRPMLREVLEHPWITANSS 263

Scoring table: BLOSUM62

Xgapop 10.0 , Xgapext 0.5 Ygapop 10.0 , Ygapext 0.5 Fgapop 6.0 , Fgapext 7.0 Delop 6.0 , Delext 7.0

Searched: 4996997 seqs, 3332346308 residues

Total number of hits satisfying chosen parameters: 99933994

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Command line parameters:

- -MODEL=frame+ p2n.model -DEV=xlp
- -Q=/abss/ABSSWEB_spool/US10601011/runat_12022006_150910_13321/app query.fasta 1
- -DB=N_Geneseq -QFMT=fastap -SUFFIX=rng -MINMATCH=0.1 -LOOPCL=0 -LOOPEXT=0
- -UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi -LIST=45
- -DOCALIGN=200 -THR_SCORE=pct -THR_MAX=100 -THR_MIN=0 -ALIGN=15 -MODE=LOCAL
- -OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=200000000 -HOST=abss05p
- -USER=US10601011_@CGN_1_1_1096_@runat_12022006_150910_13321 -NCPU=6 -ICPU=3
- -NO MMAP -NEG SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG -DEV TIMEOUT=120
- -WARN TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6 -FGAPEXT=7
- -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7

Database : N_Geneseq_21:*

- 1: geneseqn1980s:*
- 2: geneseqn1990s:*
- 3: geneseqn2000s:*
- 4: geneseqn2001as:*
- 5: geneseqn2001bs:*
- 6: geneseqn2002as:*
- 7: geneseqn2002bs:*
- 8: geneseqn2003as:*
- 9: geneseqn2003bs:*
- 10: geneseqn2003cs:*
- 11: geneseqn2003ds:*
- 12: geneseqn2004as:*

13: geneseqn2004bs:*
14: geneseqn2005s:*

용

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

SUMMARI

Res	n1+		Query				
	No.	Score		Length	DB	ID	Description
							Description
	1	1383	100.0	2005	10	ADF61839	Adf61839 Human ser
	2	1383	100.0	2070	6	ABQ55035	Abq55035 Human ova
	3	1383	100.0	2146	8	ACA03962	Aca03962 cDNA down
	4	1383	100.0	2183	13	ADT90062	Adt90062 Human gen
	5	1383	100.0	2198	2	AAT67290	Aat67290 Human Aur
	6	1383	100.0	2198	2	AAX99725	Aax99725 Human AUR
	7	1383	100.0	2237	6	AAS94997	Aas94997 Human DNA
	8	1383	100.0	2253	6	ABK64863	Abk64863 DNA encod
	9	1383	100.0	2253	8	ACC50970	Acc50970 Human bla
	10	1383	100.0	2253	10	ADG89375	Adg89375 Cancer de
	11	1383	100.0	2253	11	ADN39671	Adn39671 Cancer/an
	12	1383	100.0	2253	12	ADH59555	Adh59555 DNA seque
	13	1383	100.0	2253	12	ADM72204	Adm72204 Human TAS
	14	1383	100.0	2253	13	ADR25068	Adr25068 Breast ca
	15	1383	100.0	2253	13	ADT89948	Adt89948 Human gen
	16	1383	100.0	2253	13	ADT90311	Adt90311 Human gen
	17	1383	100.0	2253	13	ADT90511	Adt90571 Human gen
	18	1383	100.0	2253	13		
	19	1383	100.0			ADT90607	Adt90607 Human gen
				2253	13	ADT90179	Adt90179 Human gen
	20	1383	100.0	2253	13	ADU05998	Adu05998 Novel bro
	21	1383	100.0	2253	13	ADX97464	Adx97464 Pancreati
	22	1383	100.0	2253	14	ADX06368	Adx06368 Cyclin-de
	23	1383	100.0	2253	14	ADY15253	Ady15253 DNA encod
	24	1383	100.0	2253	14	ADY61799	Ady61799 Human gen
	25	1383	100.0	2283	11	ACN89131	Acn89131 Breast ca
С	26	1383	100.0	2284	5	ADL63462	Adl63462 Human ova
	27	1383	100.0	2346	13	ADR05175	Adr05175 Human GTP
	28	1383	100.0	2347	10	ABZ75005	Abz75005 Human ser
	29	1383	100.0	2347	10	ABZ75006	Abz75006 Human ser
	30	1383	100.0	2347	13	ACN40798	Acn40798 Tumour-as
	31	1363	98.6	2148	6	AAS94996	Aas94996 Human DNA
	32	1332	96.3	1209	10	ACF58068	Acf58068 Human HsA
	33	1332	96.3	1209	13	ADQ89831	Adq89831 Antagonis
	34	1332	96.3	2033	6	ABK64870	Abk64870 DNA encod
	35	1332	96.3	2033	10	ADD89968	Add89968 Human can
	36	1332	96.3	2033	12	ADK67713	Adk67713 Human mod
	37	1332	96.3	2033	13	ADR25020	Adr25020 Breast ca
	38	1332	96.3	2033	13	ADU06017	Adu06017 Novel bro
	39	1332	96.3	2033	14	ADX06366	Adx06366 Cyclin-de
	40	1332	96.3	2033	14	ADY15913	Ady15913 DNA encod
	41	1332	96.3	2033	14	ADY61794	Ady61794 Human gen
	42	1284	92.8	1500	6	ABT06285	Abt06285 Human NOV
	43	1044	75.5	1035	13	ADQ89833	Adq89833 Antagonis
	44	1044	75.5	1224	8	ACC51000	Acc51000 Human bla
	45	1044	75.5	1224	8	ABX76370	Abx76370 Lung canc

OM protein - nucleic search, using frame plus p2n model

Run on: February 14, 2006, 21:18:18; Search time 878 Seconds

(without alignments)

2477.046 Million cell updates/sec

Title: US-10-601-011-1 COPY 126 388

Perfect score: 1383

Sequence: 1 RQWALEDFEIGRPLGKGKFG.....SQRPMLREVLEHPWITANSS 263

Scoring table: BLOSUM62

Xgapop 10.0 , Xgapext 0.5 Ygapop 10.0 , Ygapext 0.5 Fgapop 6.0 , Fgapext 7.0 Delop 6.0 , Delext 7.0

Searched: 9793542 seqs, 4134689005 residues

Total number of hits satisfying chosen parameters: 19587084

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Command line parameters:

- -MODEL=frame+_p2n.model -DEV=xlh
- -Q=/abss/ABSSWEB_spool/US10601011/runat_12022006_150918 13487/app query.fasta 1
- -DB=Published_Applications NA_Main -QFMT=fastap -SUFFIX=rnpbm -MINMATCH=0.1
- -LOOPCL=0 -LOOPEXT=0 -UNITS=bits -START=1 -END=-1 -MATRIX=blosum62
- -TRANS=human40.cdi -LIST=45 -DOCALIGN=200 -THR_SCORE=pct -THR MAX=100
- -THR_MIN=0 -ALIGN=15 -MODE=LOCAL -OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0
- -MAXLEN=2000000000 -HOST=abss03h
- -USER=US10601011_@CGN_1_1 1026 @runat 12022006 150918 13487 -NCPU=6 -ICPU=3
- -NO MMAP -NEG SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG -DEV TIMEOUT=120
- -WARN TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6 -FGAPEXT=7
- -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7

Database : Published_Applications_NA Main:*

- 1: /cgn2_6/ptodata/1/pubpna/US07_PUBCOMB.seq:*
- 2: /cgn2_6/ptodata/1/pubpna/US08_PUBCOMB.seq:*
- 3: /cgn2_6/ptodata/1/pubpna/US09A PUBCOMB.seq:*
- 4: /cgn2_6/ptodata/1/pubpna/US09B_PUBCOMB.seq:*
- 5: /cgn2_6/ptodata/1/pubpna/US10A_PUBCOMB.seq:*
- 6: /cgn2_6/ptodata/1/pubpna/US10B_PUBCOMB.seq:*
- 7: /cgn2_6/ptodata/1/pubpna/US10C_PUBCOMB.seq:*
- 8: /cgn2_6/ptodata/1/pubpna/US10D_PUBCOMB.seq:*
- 9: /cgn2_6/ptodata/1/pubpna/US10E_PUBCOMB.seq:*
- 10: /cgn2_6/ptodata/1/pubpna/US11 PUBCOMB.seq:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

		•			SUMMARIES	
		*				
Result		Query				
No.	Score	Match	Length	DB	ID	Description
1		100.0	2070	6	US-10-264-049-915	Sequence 915, App
2	1383	100.0	2146		US-10-071-766-83	Sequence 83, Appl
3	1383	100.0	2198	3	US-09-012-135A-2	Sequence 2, Appli
4	1383	100.0	2198	3	US-09-784-332-2	Sequence 2, Appli
5	1383	100.0	2198	8	US-10-734-126-2	Sequence 2, Appli
6	1383	100.0	2237	6	US-10-240-965-252	Sequence 252, App
7	1383	100.0	2253	6	US-10-172-118-929	Sequence 929, App
8	1383	100.0	2253	6	US-10-388-360-323	Sequence 323, App
9	1383	100.0	2253	6	US-10-295-027-989	Sequence 989, App
10	1383	100.0	2253	7	US-10-342-887-929	Sequence 929, App
11	1383	100.0	2253	7	US-10-188-832-34	Sequence 34, Appl
12	1383	100.0	2253	8	US-10-751-736-2	Sequence 2, Appli
13	1383	100.0	2253	8	US-10-781-581-186	Sequence 186, App
14		100.0	2253	9	US-10-848-755A-115	Sequence 115, App
15		100.0	2283	5	US-10-198-846-10281	Sequence 10281, A
c 16		100.0	2284	3	US-09-814-353-21674	Sequence 21674, A
17		100.0	2347	5	US-10-209-324-1	Sequence 1, Appli
18		98.6	2148	6	US-10-240-965-251	Sequence 251, App
19		96.3	1209	9	US-10-745-237-261	Sequence 261, App
20		96.3	2033	6	US-10-172-118-881	Sequence 881, App
21		96.3	2033	7	US-10-342-887-881	Sequence 881, App
22		96.3	2033	9	US-10-848-755A-110	Sequence 110, App
23		96.3	2033	9	US-10-509-599-13	Sequence 13, Appl
24		92.8	1500	7	US-10-087-684-13	Sequence 13, Appl
25		92.8	1500	7	US-10-218-779-13	Sequence 13, Appl
26		75.5	1035	9	US-10-745-237-263	Sequence 263, App
27		75.5	1224	5	US-10-171-311-213	Sequence 213, App
28		75.5	1224	6	US-10-172-118-1004	Sequence 1004, Ap
29		75.5	1224	6	US-10-295-027-202	Sequence 202, App
30		75.5	1224	6	US-10-173-999-100	Sequence 100, App
31		75.5	1224	7	US-10-342-887-1004	Sequence 1004, Ap
32		75.5	1224	7	US-10-188-832-92	Sequence 92, Appl
33		75.5	1244	3	US-09-012-135A-1	Sequence 1, Appli
34		75.5	1244	3	US-09-784-332-1	Sequence 1, Appli Sequence 1, Appli
35		75.5	1244	8	US-10-734-126-1	Sequence 1, Appli Sequence 1, Appli
36		75.5	1244	_	US-10-335-053-258	_
37		75.4	1168	8 3		Sequence 258, App Sequence 1709, Ap
38		75.4	1168	<i>5</i>	US-09-920-300A-1709 US-10-033-528-1709	-
						Sequence 1709, Ap
39		75.4	1168	6	US-10-099-926-1709	Sequence 1709, Ap
40		75.4	1168	9	US-10-961-527-1709	Sequence 1709, Ap
41		74.7	1272	3	US-09-974-298-135	Sequence 135, App
42		74.7	1272	5	US-10-116-802-135	Sequence 135, App
43		74.7	1272	5	US-10-084-817-314	Sequence 314, App
44		74.5	1197	7	US-10-403-571-97	Sequence 97, Appl
45	1031	74.5	1923	9	US-10-764-420-1670	Sequence 1670, Ap

OM protein - protein search, using sw model

Run on: February 12, 2006, 18:58:35; Search time 43 Seconds

(without alignments)

588.489 Million cell updates/sec

Title: US-10-601-011-1_COPY_126_388

Perfect score: 1383

Sequence: 1 RQWALEDFEIGRPLGKGKFG.....SQRPMLREVLEHPWITANSS 263

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 283416 seqs, 96216763 residues

Total number of hits satisfying chosen parameters: 283416

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : PIR_80:*

1: pir1:*

2: pir2:*

3: pir3:*

4: pir4:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

	용				
	Query				
Score	Match	Length	DB	ID	Description
1374	99.3	403	2	JC5974	aurora-related kin
1292.5	93.5	395	2	JC5975	aurora-related kin
1178	85.2	389	2	S52242	protein kinase (EC
1174	84.9	407	2	S52243	p46Eg265 protein -
1025	74.1	345	2	JC4665	protein kinase (EC
933.5	67.5	282	2	H84653	probable protein k
925.5	66.9	294	2	T10690	serine/threonine-s
895.5	64.8	384	2	T41298	ser/thr protein ki
891	64.4	421	2	A56220	protein kinase (EC
890	64.4	305	2	T43221	serine/threonine-s
890	64.4	329	2	B87790	protein B0207.4 [i
858.5	62.1	288	2	T00862	probable serine/th
769	55.6	326	2	T43219	serine/threonine-s
	1374 1292.5 1178 1174 1025 933.5 925.5 895.5 891 890 890 858.5	Query Score Match 1374 99.3 1292.5 93.5 1178 85.2 1174 84.9 1025 74.1 933.5 67.5 925.5 66.9 895.5 64.8 891 64.4 890 64.4 890 64.4 890 64.4	Query Score Match Length	Query Score Match Length DB 1374 99.3 403 2 1292.5 93.5 395 2 1178 85.2 389 2 1174 84.9 407 2 1025 74.1 345 2 933.5 67.5 282 2 925.5 66.9 294 2 895.5 64.8 384 2 891 64.4 421 2 890 64.4 305 2 890 64.4 329 2 858.5 62.1 288 2	Query Score Match Length DB ID 1374 99.3 403 2 JC5974 1292.5 93.5 395 2 JC5975 1178 85.2 389 2 S52242 1174 84.9 407 2 S52243 1025 74.1 345 2 JC4665 933.5 67.5 282 2 H84653 925.5 66.9 294 2 T10690 895.5 64.8 384 2 T41298 891 64.4 421 2 A56220 890 64.4 305 2 T43221 890 64.4 329 2 B87790 858.5 62.1 288 2 T00862

14	769	55.6	343	2	D89124
15	701.5	50.7	367	2	S47923
16	505	36.5	1650	2	T18444
17	501.5	36.3	465	2	B55748
18	501.5	36.3	925	2	A55748
19	493.5	35.7	440	2	T14735
20	481	34.8	399	1	OKBYC1
21	479.5	34.7	440	2	T14736
22	477.5	34.5	441	2	C84667
23	475.5	34.4	480	2	A86427
24	466	33.7	398	1	OKBYC3
25	466	33.7	512	1	JC1446
26	464	33.6	421	2	T48202
27	464	33.6	461	2	T14822
28	458.5	33.2	445	2	T50802
29	456.5	33.0	445	2	T09903
30	452	32.7	442	2	T48203
31	452	32.7	489	2	T04862
32	450	32.5	426	2	C71408
33	448	32.4	441	2	E85362
34	447.5	32.4	1398	2	T13741
35	446.5	32.3	746	2	S62365
36	445	32.2	504	2	T10449
37	445	32.2	511	1	A56009
38	444.5	32.1	512	2	A54400
39	443.5	32.1	301	1	A40811
40	443.5	32.1	510	2	T04145
41	443	32.0	380	1	OKBYC2
42	441	31.9	425	2	S41099
43	440	31.8	481	2	JE0377
44	439	31.7	648	1	JQ1150
45	438	31.7	1245	2	D86260

protein K07C11.2 [probable protein k hypothetical prote protein kinase (EC protein kinase (EC probable serine/th protein kinase (EC probable serine/th probable protein k probable serine/th protein kinase (EC serine/threonine-s protein kinase AK2 probable serine/th serine/threonine p serine/threonine-s hypothetical prote probable serine/th probable protein k hypothetical prote hypothetical prote SNF1-related prote probable serine/th serine/threonine-s protein kinase (EC myosin-light-chain serine/threonine p protein kinase (EC protein kinase (EC p70 S6 kinase (EC protein kinase (EC protein T12C24.22

OM protein - protein search, using sw model

Run on: February 12, 2006, 18:57:06; Search time 252 Seconds

(without alignments)

736.325 Million cell updates/sec

Title: US-10-601-011-1 COPY 126 388

Perfect score: 1383

Sequence: 1 RQWALEDFEIGRPLGKGKFG.....SQRPMLREVLEHPWITANSS 263

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2166443 seqs, 705528306 residues

Total number of hits satisfying chosen parameters: 2166443

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : UniProt 05.80:*

1: uniprot_sprot:*
2: uniprot_trembl:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

		8				
Result		Query				
No.	Score	Match	Length	DB	ID	Description
1	1383	100.0	403	1	STK6_HUMAN	014965 homo sapien
2	1334.5	96.5	405	2	Q4R1K4_PIG	Q4r1k4 sus scrofa
3	1310	94.7	397	1	STK6_RAT	P59241 rattus norv
4	1305	94.4	386	2	Q8C3H8_MOUSE	Q8c3h8 mus musculu
5	1305	94.4	395	1	STK6_MOUSE	P97477 mus musculu
6	1305	94.4	395	2	Q7TNK2_MOUSE	Q7tnk2 mus musculu
7	1294	93.6	395	2	Q8BP87_MOUSE	Q8bp87 mus musculu
8	1178	85.2	408	1	STK6L_XENLA	Q91819 xenopus lae
9	1176	85.0	408	2	Q6DJK0_XENLA	Q6djk0 xenopus lae
10	1174	84.9	407	1	STK6_XENLA	Q91820 xenopus lae
11	1174	84.9	407	2	Q6INY9_XENLA	Q6iny9 xenopus lae
12	1150	83.2	347	2	Q4R9G2_MACFA	Q4r9g2 macaca fasc
13	1146	82.9	343	2	Q5QPD5_HUMAN	Q5qpd5 homo sapien
14	1113	80.5	405	2	Q6DBZ4_BRARE	Q6dbz4 brachydanio
15	1075	77.7	367	2	Q4SS89_TETNG	Q4ss89 tetraodon n

16	1056	76.4	320	2	Q6NW76_BRARE	Q6nw76 brachydanio	
17	1054	76.2	368	2	Q6GPL3_XENLA	Q6gpl3 xenopus lae	
18	1051	76.0	361	2	Q9DF70_XENLA	Q9df70 xenopus lae	
19	1047	75.7	361	2	Q8JG74_XENLA	Q8jg74 xenopus lae	
20	1047	75.7	361	2	Q6DE08_XENLA	Q6de08 xenopus lae	
21	1047	75.7	371	2	Q7ZYT9_XENLA	Q7zyt9 xenopus lae	
22	1044	75.5	344	1	AURKB_HUMAN	Q96gd4 homo sapien	
23	1031	74.5	344	2	Q7YRC7_PIG	Q7yrc7 sus scrofa	
24	1031	74.5	345	1	AURKB_MOUSE	070126 mus musculu	
25	1031	74.5	345	2	Q8C6C1_MOUSE	Q8c6c1 mus musculu	
26	1031	74.5	346	2	Q8JGS8_BRARE	Q8jgs8 brachydanio	
27	1029	74.4	343	1	AURKB_RAT	055099 rattus norv	
28	1029	74.4	343	2	Q4V8N1_RAT	Q4v8n1 rattus norv	
29	1025	74.1	344	2	Q7YRC6_BOVIN	Q7yrc6 bos taurus	
30	1017	73.5	289	2	Q6AZY8_HUMAN	Q6azy8 homo sapien	
31	1017	73.5	290	2	Q6DLZ0_HUMAN	Q6dlz0 homo sapien	
32	1017	73.5	306	2	Q5Y191_HUMAN	Q5y191 homo sapien	
33	1017	73.5	309	1	AURKC_HUMAN	Q9uqb9 homo sapien	
34	1007	72.8	270	2	Q4RTH5_TETNG	Q4rth5 tetraodon n	
35	963	69.6	276	2	Q6P209_MOUSE	Q6p209 mus musculu	
36	963	69.6	282	1	AURKC_MOUSE	088445 mus musculu	
37	935.5	67.6	292	2	Q5SNH4_ORYSA	Q5snh4 oryza sativ	
38	933.5	67.5	282	2	082309_ARATH	082309 arabidopsis	
39	933.5	67.5	288	2	Q683C9_ARATH	Q683c9 arabidopsis	
40	926	67.0	432	2	Q4R1K7_ORYSA	Q4r1k7 oryza sativ	
41	925.5	66.9	294	2	Q9M077_ARATH	Q9m077 arabidopsis	
42	924.5	66.8	294	2	Q8LBX4_ARATH	Q8lbx4 arabidopsis	
43	913	66.0	404	2	Q7PY85_ANOGA	Q7py85 anopheles g	
44	901	65.1	302	2	Q61XD3_CAEBR	Q61xd3 caenorhabdi	
45	895.5	64.8	355	1	ARK1_SCHPO	059790 schizosacch	

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